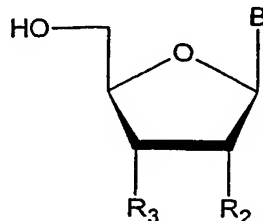


ClaimsREPLACED BY  
ART 34 AMDT

1. A method for preparing an oligonucleotide comprising the steps of

a) providing a 3'-protected compound having the formula:



5 wherein

B is a heterocyclic base

R<sub>2</sub> is H, a protected 2'-hydroxyl group, F, a protected amino group, an O-alkyl group, an O-substituted alkyl, a substituted alkylamino or a C4'-O2' methylen linkage

10 R<sub>3</sub> is OR'<sub>3</sub>, NHR'<sub>3</sub>, NR''<sub>3</sub>R'''<sub>3</sub>, a 3'-protected nucleotide or a 3'-protected oligonucleotide,

R'<sub>3</sub> is a hydroxyl protecting group,

R''<sub>3</sub>, R'''<sub>3</sub> are independently an amine protecting group,

15 b) reacting said compound with a nucleotide derivative having a 5'-protection group in the presence of a solid supported activator to give an elongated oligonucleotide with a P(III)-internucleotide bond

c) optionally processing the elongated oligonucleotide with a P(III)-internucleotide bond by either or both of steps c1) and c2) in any sequence

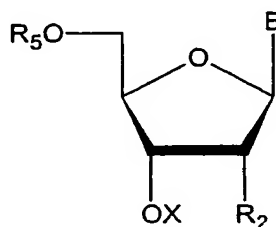
20 c1) capping preferably by reacting with a solid supported capping agent

c2) oxidizing preferably by reacting the oligonucleotide with a solid supported oxidizing reagent

d) removing the 5'-protection group.

25 2. The method of claim 1, wherein the step d) is effected by treatment with a solid supported agent or removing the 5'-protection group with a removal agent followed by addition of a solid supported scavenger or followed by extraction.

3. The method of claim 1 or 2, wherein the nucleotide derivative having a 5'-protection group of step b) has the following formula:



wherein

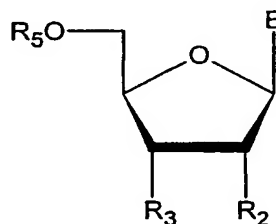
X is a P(III)-function

B is a heterocyclic base

R<sub>2</sub> is H, a protected 2'-hydroxyl group, F, a protected amino group, an O-alkyl group, an O-substituted alkyl, a substituted alkylamino or a C4'-O2' methylen linkage

R<sub>5</sub> is a hydroxyl protecting group, a 5'-protected nucleotide or a 5'-protected oligonucleotide.

4. A method for preparing an oligonucleotide comprising the steps of  
a) providing a 5'-protected compound having the formula:



wherein

B is a heterocyclic base

R<sub>2</sub> is H, a protected 2'-hydroxyl group, F, a protected amino group, an O-alkyl group, an O-substituted alkyl, a substituted alkylamino or a C4'-O2' methylen linkage

R<sub>3</sub> is OH, NH<sub>2</sub>

R<sub>5</sub> is a hydroxyl protecting group, a 5'-protected nucleotide or a 5'-protected oligonucleotide

- b) reacting said compound with a nucleotide derivative having a 3'-

protection group in the presence of a solid supported activator to give an elongated oligonucleotide with a P(III)-internucleotide bond

c) optionally processing the elongated oligonucleotide with a P(III)-internucleotide bond by either or both of steps c1) and c2) in any sequence

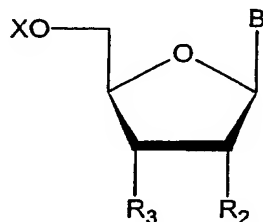
c1) capping, preferably by reacting with a solid supported capping agent

c2) oxidizing, preferably by reacting the oligonucleotide with a solid supported oxidizing reagent

d) removing the 3'-protection group.

5. The method of claim 4, wherein step d) is effected by treatment with a solid supported agent or removing the 3'-protection group with a removal agent followed by addition of a solid supported scavenger or followed by extraction.

6. The method of claim 4 or 5, wherein the nucleotide derivative having a 3'-protection group has the following formula:



wherein

X is a P(III)-function

B is a heterocyclic base

R<sub>2</sub> is H, a protected 2'-hydroxyl group, F, a protected amino group, an O-alkyl group, an O-substituted alkyl, a substituted alkylamino or a C4'-O2' methylen linkage

R<sub>3</sub> = OR'<sub>3</sub>, NHR''<sub>3</sub>, NR'''<sub>3</sub>R'''', a 3'-protected nucleotide or a 3'-protected oligonucleotide,

R'<sub>3</sub> is a hydroxyl protecting group,

R''<sub>3</sub>, R'''<sub>3</sub> are independently an amine protecting group,

R'<sub>3</sub> is a hydroxyl protecting group, a 3'-protected nucleotide or a 3'-protected oligonucleotide

7. The method of any one of claims 1 to 5, comprising the further step of  
e) repeating steps a) to d) at least once.
8. The method of any one of claims 1 to 6, wherein the nucleotide derivative of  
step b) is a phosphoramidite or a H-phosphonate.
- 5 9. The method of any one of steps 1 to 8, wherein the solid supported activator  
of step b) is selected from the group consisting of a solid support bearing a  
pyridinium salt, a cation exchange solid support with an optionally substi-  
tuted pyridinium, a cation exchange solid support with an optionally substi-  
tuted imidazolium salt, a solid support bearing an optionally substituted azole  
10 (imidazol, triazole, tetrazole), a salt of a weak base anion exchange resin  
with a strong acid, a weak cation exchange resin (carboxylic) in its proto-  
nated form, a solid support bearing an optionally substituted phenol, a solid  
support bearing a carboxylic acid chloride/bromide, a sulfonic acid chlo-  
ride/bromide, a chloroformate, a bromoformate, a chlorosulfite, a bromosul-  
15 fite, a phosphorochloridate, a phosphorbromidate and a solid support bound  
carbodiimide.
10. The method of any one of claims 1 to 9, wherein the solid supported oxidiz-  
ing reagent is selected from the group consisting of solid supported perio-  
dates, permanganates, osmium tetroxides, dichromates, hydroperoxides,  
20 substituted alkylamine oxides, percarboxylic acid and persulfonic acid.
11. The method of any one of claims 1 to 10, wherein the oxidizing is a sulfuri-  
zation.
12. The method of claim 11, wherein the solid supported oxidizing reagent is  
selected from the group consisting of a solid supported tetrathionate, a solid  
25 supported alkyl or aryl sulfonyl disulfide, a solid supported optionally substi-  
tuted dibenzoyl tetrasulfide, a solid supported bis(alkoxythio-  
carbonyl)tetrasulfide, a solid supported optionally substituted phenylacetyl  
disulfide, a solid supported N-[(alkyl or aryl)sulfanyl] alkyl or aryl substituted  
succinimide and a solid supported (2-pyridinyldithio) alkyl or aryl.
- 30 13. The method of any one of claims 1 to 12, wherein the solid supported cap-  
ping agent is a solid supported activated acid, preferably a carboxylic acid  
chloride, carboxylic acid bromide, azolide, substituted azolide, anhydride or  
chloroformate or phosphorochloridate, or a solid supported phosphoramidite,  
or a solid supported H-phosphonate monoester.

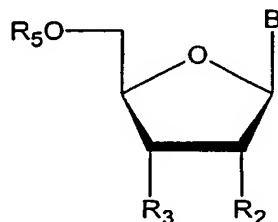
14. The method of any one of claims 1 to 13, wherein the 5'-protection is a dimethoxytrityl group (DMTr) or a monomethoxytrityl group (MMTr) and the solid supported agent of step d) is an cationic ion exchanger resin in the H<sup>+</sup> form or solid supported ceric ammonium nitrate.

15. The method of any one of claims 1 to 14, wherein the 3'-protection is a silyl group and the solid supported agent of step d) is an anionic ion exchanger resin in the F-form or the 3'-protection is levulinic acid and the solid supported agent of step d) is a solid supported hydrazine or a solid supported hydrazinium.

16. Use of a solid supported sulfurization agent consisting of solid supported amine and a tetrathionate having the formula S<sub>4</sub>O<sub>6</sub> or a cyanoethylthiosulfate (NC-CH<sub>2</sub>-CH<sub>2</sub>-S-SO<sub>3</sub><sup>-</sup>) for sulfurization of oligonucleotides.

17. A method for preparing an oligonucleotide comprising the steps of

a) providing a compound having the formula:



wherein

B is a heterocyclic base

R<sub>2</sub> is H, a protected 2'-hydroxyl group, F, a protected amino group, an O-alkyl group, an O-substituted alkyl, a substituted alkylamino or a C4'-O2' methylene linkage

and

R<sub>3</sub> is OR'<sub>3</sub>, NHR''<sub>3</sub>, NR'''<sub>3</sub>R''''<sub>3</sub>,

a protected nucleotide or a protected oligonucleotide and R<sub>5</sub> is a P(III) function

R'<sub>3</sub> is a hydroxyl protecting group,

R''<sub>3</sub>, R'''<sub>3</sub> are independently an amine protecting group,

or

R<sub>5</sub> is a hydroxyl protecting group, a protected nucleotide or a protected oligonucleotide and R<sub>3</sub> is a P(III) function

- b) reacting said compound with a nucleotide derivative having a 3' or 5'-free OH-group in the presence of a solid supported activator to give an elongated oligonucleotide with a P(III)-internucleotide bond
- 5 c) optionally processing the elongated oligonucleotide with a P(III)-internucleotide bond by either or both of steps c1) and c2) in any sequence
- c1) capping by reacting with a solid supported capping agent
- c2) oxidizing by reacting the oligonucleotide with a solid supported oxidizing reagent
- 10 d) removing the 3' or 5'-protection group.

REPLACED BY  
ART 34 AMDT